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Diminished alpha lateralization during Working Memory but not during attentional cueing in older adults

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25

Abstract

Aging has been associated with declined performance in tasks that rely on working 26 memory (WM). Because attention and WM are tightly coupled, declined performance on a 27 28 WM task in older adults could be due to deficits in attention, memory capacity, or both. We used alpha (8-14 Hz) power modulations as an index to assess how changes in attention and 29 memory capacity contribute to decreased WM performance in older adults. We recorded the 30 magnetoencephalogram in healthy older (60–76 years) and younger adults (18–28 years) 31 while they performed a lateralized WM task. At matched difficulty, older adults showed 32 significantly lower memory-spans than younger adults. Alpha lateralization during retention 33 was nearly absent in older adults due to a bilateral reduction of alpha power. By contrast, in 34 35 younger adults alpha power was reduced only contralateral to the attended hemifield. 36 Surprisingly, during the cue interval, both groups showed equal alpha lateralization. The preserved alpha lateralization during attentional cueing, and lack thereof during retention, 37 suggests that reduced WM performance in older adults is due to deficits in WM-related 38 processes, not deficits in attentional orienting, and that a compensatory mechanism in aging 39 that permits significant residual WM performance in the absence of alpha lateralization. 40 Keywords: Attention, Healthy aging, MEG, Oscillations 41

42 Growing old is characterized by general cognitive slowing and decline (for a review, see e.g. Hedden and Gabrieli, 2004), which often involves working memory (WM) deficits 43 (Cabeza et al. 2002; Park et al. 2002; Bopp and Verhaeghen 2005; Sander et al. 2011; Murre 44 45 et al. 2013). WM refers to the ability to briefly store information for later use (D'Esposito 2007), and is crucial for many types of cognition. WM includes encoding, retention, and 46 recollection or recognition phases. It has limited capacity, and most individuals can only store 47 three to four items (Luck and Vogel 1997, 2013; Cowan 2000; Vogel et al. 2001). This 48 limited capacity requires efficient use of resources, and thus WM benefits from an attentional 49 50 filter that prevents encoding of irrelevant stimuli, thereby limiting encoding to the relevant stimuli. As a result, attention and WM are closely interrelated, and declined WM 51 52 performance in older adults has indeed frequently been linked to deficits in selective attention 53 (Vogel et al. 2005; Gazzaley et al. 2008; Jost et al. 2011; Sander et al. 2011; McNab et al. 2015). 54

A striking finding when using tasks that require lateralized covert attention, such as the Delayed Match-to-Sample (DMS) task in Vogel and Machizawa (2004), is that alpha power (8 – 14 Hz) in the hemisphere contralateral to a relevant stimulus is lower than in the hemisphere contralateral to an irrelevant stimulus (Worden et al. 2000; Thut 2006; Sauseng et al. 2009; Händel et al. 2011). This observation gave rise to the current understanding of alpha oscillations as reflecting the active suppression of both encoding and maintenance of irrelevant stimuli in WM.

These experiments have thus far been performed almost exclusively in younger adults and it remains unclear to what extent these findings can be generalized to older adults, and to what extent attention and WM deficits associated with aging may be reflected by changes in the bilateral distribution of alpha. To our knowledge, there have been only three earlier studies that investigated aging and alpha lateralization during a WM task. Using EEG, Sander

67 et al. (2012b) found that older adults showed lateralized alpha power during a 1000 ms retention interval for medium loads, but not for high memory loads. Younger adults on the 68 other hand showed lateralized alpha power under both high and medium loads. In another 69 70 EEG study using a cued target discrimination paradigm, Hong et al. (2015) found that unlike younger adults, older adults did not show lateralized alpha oscillations during spatial 71 72 attention in a 1000 ms interval following a 200 ms directional cue. Recently, measuring MEG in older adults, Mok et al. (2016) found that older adults retain the ability to orient attention 73 within WM, as evidenced by alpha lateralization in response to a so called 'retro-cue', a cue 74 75 that turned on after bilateral stimulus presentation. Taken together, these results point to age related changes in alpha lateralization. However, it is as yet unclear to what extent these 76 77 observations generalise to other tasks, and moreover whether they are specific to either 78 spatial attention or WM-related processes.

Here we combine a lateralized DMS task with MEG to investigate the differences in 79 alpha lateralization between younger and older adults. We recorded alpha power during a 80 81 prolonged cueing interval and a subsequent retention interval of equal length. This paradigm allowed us to investigate whether age related differences in alpha modulation were specific to 82 either attentional cueing or WM retention, or were a general feature of both processes. A key 83 element of our study was that the directional cue remained visible throughout each trial, 84 eliminating the need to keep the directional cue in memory. The cueing interval in our study 85 86 thus represented a period of spatial attention without WM contribution, and any changes in alpha modulation in this interval could be interpreted in terms of spatial attention. The WM 87 retention interval combines attentional processes with maintenance of WM content. It is thus 88 difficult to separate WM from its associated attentional processes (Gazzaley and Nobre 2012). 89 However, our design offers a step forward in permitting the separate probing of attention 90

- 91 during the spatial cueing interval and of WM and its associated processes during the retention
- 92 interval.

Materials and Methods

94 **Participants**

Forty-six older adults were recruited via advertising at an on-campus education center 95 96 for older adults, by advertisement in the Donders Institute's participant waiting room, and via the first author's network of colleagues. Of these forty-six adults, three participants did not 97 pass an initial screening on performance (less than 60% accuracy on the lowest load) and/or 98 MRI/MEG compatibility (due to metal implants). Four more participants were excluded from 99 100 the experiment after the MRI measurement due to claustrophobic complaints and the 101 discovery of implants that had not been reported to the researchers. Five more participants were excluded after the MEG session, due to MEG (DSQ) electronics errors, excessive head 102 103 motion and muscle artifacts.

104 Data for the remaining thirty-four participants (21 men, 13 women), 60-76 years old (M = 65.8 years), was fully analyzed. All participants reported to be right-handed and had normal 105 or corrected-to-normal vision, as assessed with a Landolt C chart. Participants with glasses 106 were given MEG compatible lenses, such that they were able to read the Landolt C chart 107 equally well with the MEG compatible lenses as with their prescription glasses. We did not 108 test visual acuity in participants who used contact lenses or who did not wear glasses or 109 lenses. All participants were screened with a Dutch test resembling the Mini-Mental State 110 Examination, known as the "Cognitieve Screening Test" (De Graaf and Deelman 1991). All 111 112 participants scored normally on the screening test, indicating the absence of any major neuropsychiatric disorders. 113

We compared task performance and MEG data from the group of older adults to parallel data acquired in younger adults during an earlier experiment with the same task (Lozano-Soldevilla et al. 2014). In that experiment, the influence of GABA on visual gamma and alpha oscillations was investigated by exposing participants to an experimental and a

118	placebo session. For the current experiment, we used the placebo sessions of 25 participants
119	(12 men, 13 women), aged 18-28 years ($M = 22.4$ years) as a control, after eliminating 7 of
120	32 recruited participants according to the same criteria used for the older participants (for
121	details, see the supplementary material of Lozano-Soldevilla et al., 2014).
122	
123	Figure 1 near here
124	
125	Task and stimuli
126	For the behavioral tests, stimuli were presented on a 24" BenQ LED TFT-monitor
127	(1920x1080 px, 120 Hz refresh rate). For the MEG recordings we used an EIKI LCD
128	projector (60 Hz) projecting stimuli onto the back of a translucent screen via two mirrors.
129	We used an adapted version of a classic lateralized delayed match-to-sample task (cf.
130	Vogel and Machizawa, 2004). Participants have to decide whether an array of stimuli
131	presented in an attended hemifield is identical to a remembered array, while ignoring stimuli
132	in the other hemifield. In our version of the task, participants had to merely report the
133	presence or absence of a color change in one of the stimuli in the attended hemifield (Figure
134	1).
135	Each trial started with a fixation cross, followed by a 1500 millisecond cue period,
136	during which participants received an arrow cue that pointed towards the hemifield in which
137	the relevant parts of the stimulus arrays would be presented. This cue remained visible
138	throughout the trial. After an initial attentional cue period, a bilateral sample array of colored
139	squares was presented for 100 milliseconds, and participants had to memorize the colors of
140	the squares in the cued hemifield. The other (distractor) squares were irrelevant to the task.
141	After a retention period of 1500 ms in which only the cue was visible, a memory probe array
142	appeared and was presented until participants responded, up to a maximum response time of

143 2000 ms. In the attended hemifield, the probe stimuli either matched those in the sample array (50% of trials), or differed by the color of one square. Independent of the attended 144 hemifield, the irrelevant side of the probe array also differed from the sample array by one 145 146 square in half of the trials. Participants were instructed to ignore changes on the irrelevant side and only report changes in color on the relevant side. Responses were made by pressing 147 either a button indicating 'no change in colors', or another button indicating 'change in 148 colors', using the right index and middle finger. The mapping of the two response buttons to 149 change or no-change responses was randomized across participants. 150

151

152 Experimental procedure

The older participants were invited to two sessions. In the first session the procedure 153 154 was explained to the participant, and they could then opt out of the experiment or consent to participate. After giving informed consent, a screening took place which consisted of the CST, 155 and a final check on MEG and MRI eligibility. Similar to the procedure in Lozano-Soldevilla 156 157 et al. (2014), the experimenter then explained the task and participants completed 8-16 practice trials on a computer in a private cubicle with dimmed lights, in order to familiarize 158 themselves with the task. Head position was not restrained, although participants were placed 159 roughly 60 cm from the screen. These practice trials were presented with only one square in 160 each hemifield ('load 1'). After participants confirmed that they understood and were able to 161 162 perform the task, they completed 144 trials with loads of 2, 3, and 4 squares per hemifield (48 trials per load). This procedure both trained the participant and allowed us to adjust the 163 difficulty level for each participant individually before MEG acquisition. The load-condition 164 in which a participant performed with accuracy closest to 75% was selected as the load-165 condition that would be presented during the MEG session. Thus the difficulty of the task 166 was matched for all participants. In the first session, participants also underwent a structural 167

MRI scan (T1 weighted imaging, see next section). The total duration of the first session was1.5 h.

In the second session, which was always separated from the first by at least 5 days, 170 participants returned for MEG acquisition. After arriving, participants were asked whether 171 they still wanted to participate in the experiment. If so, they were again given 8-16 practice 172 trials (load 1) to refresh their memory of the task. After that, they were prepared for MEG 173 measurement. Participants with glasses received MEG compatible glasses following the 174 procedure for vision correction outlined above. EOG and ECG electrodes were placed, and 175 the participant was guided to the MEG system. Participants then completed 4 blocks of 100 176 trials of the task with the load that was selected for them based on session 1. Preparation took 177 1 hour, and the MEG acquisition was limited to 1 hour. 178

Younger adults completed the MEG acquisition session three times with at least 4 days in between, where in each session a different dosage of drug was administered, 1.5 mg, 0.5 mg and placebo control (for details, see Lozano-Soldevilla et al., 2014). Here we used the recordings from the placebo control condition.

183

184 MRI acquisition

185 T1-weighted images were acquired on a 1.5T Siemens Magnetom Avanto MRI system 186 (Siemens Healthcare, Erlangen, Germany). TR, TE, and TI were set to 2300 ms, 2.95 ms, and 187 850 ms, respectively. A flip angle of 15° was used, and 192 saggital slices were taken. The 188 purpose of these MRI scans was to screen for any brain abnormalities, and to retain the 189 possibility of conducting source analysis for future work.

190

191 MEG acquisition

Brain activity was measured using a 275 axial gradiometer MEG system (VSM

193 MedTech/CTF MEG, Coquitlam, Canada), with a sampling rate of 1200 Hz and a built-in

194 low-pass anti-aliasing filter with a cutoff at 300 Hz. Eye movements and blinks were

195 monitored using bipolar electrodes, applied above and below the left eye (vertical EOG), and

between the bilateral temples and outer canthi (horizontal EOG). To measure the heartbeat,

197 bilateral electrodes were applied above the right clavicle and below the left side ribs.

198 Impedance was kept below 10 k Ω for all applied electrodes.

Once inside the MEG helmet, participants were instructed to rest their head against the 199 200 back of the MEG helmet, to alleviate tension on the neck muscles and to gain optimal signal from posterior brain sites. To track the position of the head inside the MEG helmet, we used 201 202 three head coils placed at anatomical landmarks (nasion and both ear canals). Using a real-203 time head localizer (Stolk et al. 2013), we could track the position of the head relative to the MEG helmet. The position of a participant in the first few trials was saved as a template for 204 the rest of the recording. If a participant's head position deviated from the template beyond a 205 206 threshold of 5 mm in any direction, the measurement was paused and the participant was guided back into his or her original position. 207

208

209 Data analysis

210 <u>Behavior analysis</u>

Task performance was assessed by computing accuracy (correct responses divided by total responses). Response bias (*c*) and *d'* were also computed, using the formulas below (cf. Hautus, 1995):

$$d' = \varphi^{-1}(h+0.5) - \varphi^{-1}(f+1)$$
$$c = \frac{-(\varphi^{-1}(h+0.5) - \varphi^{-1}(f+1))}{2}$$

With *h* being the hit rate, *f* the false alarm rate, and φ^{-1} converting probabilities into zscores. *K*_{span} is a classic measure of memory span, we calculated it using Pashler's formula (Pashler 1988).

$$K_{span} = N\left(\frac{h-f}{1-f}\right)$$

This formula takes into account the memory load by multiplying the ratio with load factor N. 218

219 <u>MEG analysis</u>

220 The MEG data was analyzed using FieldTrip, an open-source toolbox (Oostenveld et al. 2011). All recordings were down-sampled to 600 Hz, and low-pass filtered at 200 Hz. The 221 continuous data was segmented into trials that started 2 s before array onset, and ended 3 s 222 after array onset (total trial length: 5 seconds). Line noise was eliminated by fitting sine and 223 cosine functions at 50, 100, and 150 Hz and subsequently subtracting these estimated 224 225 components. Trial offset was compensated by subtracting the mean. Trials were visually inspected for artifacts caused by, among other sources, muscle 226 contractions, head movement, and saccades. If such artifacts were present in a trial, the entire 227 trial was excluded from analysis. Trials without any behavioral response and trials with eye-228 229 blinks near array onset and probe onset (±500 ms) were also removed, to ensure that participants actually saw the to-be-encoded array. Eye-blink artifacts at other time-points in 230 231 each trial were identified by visually inspecting the results of an independent component

analysis (ICA; Jung et al., 2000). The same method was applied to identify fields detected by

the MEG sensors as a result of the electric activity of the heart. The MEG signal was

subsequently reconstructed from all components excluding the blink- and heart-related field

components, thus removing those from the signal.

For easier interpretation of power measurements, we created synthetic planar gradientsby comparing field gradients between horizontally and vertically adjacent axial gradiometers

238 separately, yielding two vectors per gradiometer (Bastiaansen and Knösche 2000). A time-239 frequency analysis was conducted on these vectors, before combining them by vector 240 summation. Time-frequency representations (TFR) of power were calculated by sliding a 241 time window over each trial in steps of 5 ms. Time window length was set per frequency to 242 fit 6 cycles ($\Delta t = 6/f$). Frequencies were assessed from 2 to 40 Hz in 1 Hz steps. TFRs were 243 then averaged across correct trials for each participant.

From the resulting average TFRs for correct trials the power modulation index (PMI)was computed, using the following formula:

$$PMI = \frac{(P_{left} - P_{right})}{(P_{left} + P_{right})}$$

where P_{left} is the power of a given frequency band in the 'attend left' condition and P_{right} the power of that band in the 'attend right' condition. Positive PMI values indicated that power was higher when attending left of the fixation compared to attending right, whereas negative values indicate the opposite. Thus, according to the hypothesis that higher alpha power occurs contralateral to a to-be-ignored hemifield, positive PMI values should appear in the left hemisphere ($P_{left} > P_{right}$), and negative PMI values should appear in the right hemisphere ($P_{right} > P_{left}$).

253

254 <u>Statistical Analysis</u>

In the behavioral data, group effects were tested using a two-sided independent samples t-test, with age-group as the between-group factor and a behavioral parameter (e.g. accuracy) as dependent variable. To assess functional brain differences in alpha power between the two age groups, the analysis was constrained to those sensors that were sensitive to the experimental manipulation of attention ('attend left' versus 'attend right'). To select these sensors of interest, a cluster-based nonparametric permutation test was used (Maris and Oostenveld 2007), which controls for multiple comparisons over sensors. TFRs of all 'attend

262 left' correct trials were pooled together (ignoring Age-group labels), as were the TFRs of all 'attend right' correct trials. To identify the sensors that most reliably distinguished between 263 the two attention conditions, without any contribution from WM-related processes, we used a 264 time-window from the cue interval (-1 - -0.1 s before array onset). First, a test statistic was 265 calculated for each sensor, based on a paired samples t-test with attention condition (attend 266 left versus attend right) as independent variable, and alpha power (8 - 14 Hz) as the 267 dependent variable. Sensors that were significant with p < 0.025 (two-sided t-test) were 268 clustered according to spatial adjacency. To be considered a cluster, at least three significant 269 270 adjacent sensors were required. For each cluster, t-statistics were summed. The cluster with the largest summed value was the cluster-based test statistic. 271

To test the statistical significance of the identified cluster, we applied a permutation test. 272 273 We obtained a cluster-based test statistic distribution by permuting the independent variable labels and recalculated the power differences 20000 times. At each permutation, we applied 274 the clustering algorithm, and the cluster with the largest sum of t-statistics entered the test 275 276 statistic distribution. The actual cluster-based t-statistic determined from empirical (nonpermuted) data was then compared to the distribution of permuted cluster-based t-statistics. A 277 p-value was estimated by calculating the proportion of t-statistics higher than the empirical t-278 statistic, and that p-value was then compared to the critical alpha-level of 0.05. In other words, 279 if the empirical cluster-based t-statistic fell outside of the 95% confidence interval, the null 280 281 hypothesis that the two labels were interchangeable was rejected.

The resulting significant clusters of sensors were used to compare the PMI for the two age groups. To summarize the positive and negative modulations in the left and right hemisphere, a combined PMI (cPMI) measure was created by considering the average PMI of the right hemisphere and subtracting it from the average PMI of the left hemisphere. Positive values of the resulting cPMI indicate effective modulation in the hypothesized direction. The

- two age groups were compared using a Repeated Measures ANOVA, with Interval (cue
- 288 interval vs. retention interval) as a within-subject factor, Age group (young adults vs. older
- adults) as a between-subjects factor, and cPMI value as the dependent variable.

Results

290

291 Behavioral results

292 *Memory load adjustment*

293 In the first behavioral session, we performed an experiment aimed at selecting a WM load that allowed older participants to reach the same accuracy as the younger adult control 294 group. For each older participant, we aimed to find a load setting at which accuracy was near 295 75%. To this end, we followed the same procedure as Lozano-Soldevilla et al. (2014), which 296 is outlined in the Method section. Behavioral results of the first session are summarized in 297 298 Table 1. Note that younger adults were tested up to load 6. Older adults were only tested up to load 4, as we did not expect high performance at load 5 and 6 and wished to avoid 299 300 frustrating the participants. There was a significant difference in accuracy between the two 301 age groups for load 3 (t(56)=2.43, p=0.019) and load 4 (t(56)=2.86, p=0.006). At load 2, no significant difference in accuracy was found (t(34,97) = 0.09, p = 0.93). The load that was 302 selected for each individual differed significantly between groups (t(30.35)=4.05, p=0.000), 303 with younger adults able to perform near 75% accuracy with higher loads (M = 4.12, SD =304 1.30) than older adults (M = 3.00, SD = 0.55). 305

306

307 Accuracy and reaction times

308 In the second session, participants completed the same DMS task with the individually

adjusted load. Accuracies of younger adults (M = 76%, SD = 8.2) and older adults (M =

310 80%, SD = 8.3) did not differ significantly (t(57) = -1.69, p = 0.097). The memory span

scores (Pashler's K) differed significantly between the two groups (t(39.05) = 2.71, p = 0.01),

with younger adults (M = 2.38, SD = 0.62) having higher K_{spans} than older adults (M =

313 2.00, SD = 0.41), reflecting successful performance under a higher load in younger adults.

Older adults (M = 0.97 s, SD = 0.13 s) had significantly slower reaction times (t(57) = 5.32, p

315 = 0.000) than younger adults (M = 0.76 s, SD = 0.16 s). However, a test of Spearman's rank 316 correlation between reaction times and alpha lateralization revealed no significant 317 correlations in the cue or retention intervals, in either younger or older adults (four tests, all r318 < 0.16, all p > 0.4). There were no significant differences in d'(t(56) = 1.873, p = 0.066) or 319 criterion (t(56) = -0.551, p = 0.584), indicating no age differences in sensitivity or response 320 bias (note that one younger participant could not be included in this analysis, because the 321 data-file was corrupted and single-trial performance was lost).

322

323 Suppression of distractors

We were interested in testing whether older adults correctly oriented attention in this 324 task. Therefore we tested whether they were specifically more prone to respond to stimuli 325 from the uncued hemifield. We coded trials according to whether there was a change in the 326 attended side (A^{C}) or whether there was no change (A^{NC}) , and according to whether a change 327 occurred in the unattended side or not (U^C or U^{NC}). To test whether older adults were 328 encoding both hemifields of the array, we compared participant's rate of reporting a change 329 when one occurred solely on the unattended side (A^{NC}/U^{C}) with the response rate when no 330 change occurred in either hemifield (A^{NC}/U^{NC}). We found no significant difference (paired t-331 test, t(33) = 1.30, p = 0.20) in older adults between A^{NC}/U^C trials (M = 14.8%, SD = 14.0%332 reported change) and A^{NC}/U^{NC} trials (M = 12.6%, SD = 9.9% reported change). There was 333 however a significant difference (paired t-test, t(23) = 2.60, p = 0.02) in younger adults 334 between A^{NC}/U^{C} trials (M = 16.9%, SD = 10.9% reported change) and A^{NC}/U^{NC} trials (M =335 14.2%, SD = 8.8% reported change). From this, one might conclude that younger adults were 336 more likely to respond to uncued stimuli. However when we calculated the distraction cost as 337 the contrast between those two rates for each individual $(A^{NC}/U^{C} - A^{NC}/U^{NC})$ there was no 338 significant difference (independent sample t-test, p = 0.79) between the older adults (M =339

340	2.1%, $SD = 9.5\%$) and the younger adults ($M = 2.7\%$, $SD = 5.1\%$). We also tested for
341	distractor benefit in trials where a change occurred in both sides compared to trials in which a
342	change occurred only on the attended side ($A^{C}/U^{C} - A^{C}/U^{NC}$). Response rate for A^{C}/U^{C} was
343	significantly higher than for A^{C}/U^{NC} in both young adults ($t(23) = 4.96$, $p = 0.000$) and older
344	adults ($t(33) = 2.93$, $p = 0.006$), with older adults reporting a change 3.8% ($SD = 12.9\%$)
345	more often, and young adults 5.5% (<i>SD</i> = 5.5%) more often. As before, when we tested for
346	differences in the individual subjects' contrast there was no significant difference between
347	age groups (independent sample t-test, $p = 0.33$).
348	Finally, we tested for the effect of distraction on reaction times. Although older adults
349	were slower than younger adults, they were not significantly slower (paired t-test, $p = 0.72$)
350	for U^{C} trials ($M = 0.96$ s, $SD = 0.14$ s) than for U^{NC} trials ($M = 0.97$ s, $SD = 0.13$ s). For
351	younger adults, there was no significant difference either (paired t-test, $p = 0.35$). Taken
352	together, these findings do not support the possibility that the reduced alpha lateralization in
353	older adults during WM is due to a failure to orient attention or greater interference from the
354	distractors in older adults.
355	
356	Figure 2 near here
357	
358	MEG results
359	Sensor selection
360	Figure 2A shows the results of the sensor selection. Positive values (red) indicate that

alpha power was greater in the 'attend left' condition than in the 'attend right' condition,
while negative values (blue) indicate the opposite. The cluster-based permutation test on the
grand average (all subjects combined) of normalized alpha power in the cue interval revealed
two clusters that differed significantly between the 'attend left' and 'attend right' conditions.

A significant (p = 0.004) positive cluster of 68 sensors was found over the left posterior hemisphere, and a significant (p = 0.02) negative cluster of 37 sensors was found over the right posterior hemisphere (Figure 2A, bold dots). In order to prevent a bias in sensitivity between hemispheres due to differing amounts of sensors, we selected only those sensors that were symmetrically significant in both clusters, resulting in 35 sensors per hemisphere (Figure 2A, bold black dots).

371

372 Alpha modulation and lateralization

373 Average TFRs belonging to the respective clusters during correct trials are shown in Figure 2B (young adults) and 2C (older adults). It was apparent from the TFRs that alpha 374 power modulation within the clusters was roughly similar for younger and older adults in the 375 cue interval (-1.5 s - 0 s). However, in the retention interval there was a striking difference 376 between the age groups; in younger adults alpha modulation was higher than during the cue 377 interval, whereas in older adults modulation was nearly absent. Figure 2D shows the same 378 379 data in another format, to emphasize the strong alpha power modulation during the retention interval in both hemispheres in younger adults, and the absence of such modulations in the 380 older group. In contrast, in the preceding cue interval there appeared to be no difference 381 between the age groups. 382

383

384 ------ Figure 3 near here ------

385

To quantitatively investigate these observations, we calculated combined PMI (cPMI) values by subtracting values of the negative cluster from values of the positive cluster. The cPMI values are shown in Figure 3, averaged per age group and interval. The data show similar cPMI values between younger and older adults in the cue interval, while in the

390 retention interval cPMI was clearly higher for younger adults. These observations were tested by conducting a Repeated Measures (RM) ANOVA, which revealed a significant main effect 391 of Interval (F(1,57) = 6.523, p = 0.013), with the cue interval cPMI being lower (M = 0.04, 392 SD = 0.05) than the retention interval cPMI (M = 0.06, SD = 0.08). The main effect of Age 393 group was also significant (F(1,57) = 16.943, p = 0.000), with younger adults showing higher 394 cPMI (M = 0.076, SD = 0.069) than older adults (M = 0.026, SD = 0.045). The cPMI 395 similarity in the cue interval and the cPMI difference in the retention interval resulted in a 396 significant interaction between Interval and Age group (F(1,57) = 21.15, p = 0.000). Post-hoc 397 t-tests confirmed that there was no significant difference (t(57) = 0.684, p = 0.497) between 398 the age groups during the cue interval. However, there was a highly significant difference 399 400 (t(31.50) = 4.641, p = 0.000) between younger adults (M = 0.110, SD = 0.094) and older 401 adults (M = 0.016, SD = 0.043) during the retention interval.

To exclude the possibility that the diminished alpha lateralization was due to older 402 adults making more eye-movements, we compared the rectified horizontal EOG traces during 403 404 the retention interval between young and older adults. There was no significant difference (independent samples t-test, t(54) = -0.65, p = 0.519) between the traces, although on visual 405 inspection of the traces, older adults did seem to move their eyes slightly farther. In order to 406 confidently exclude eye-movements as the cause of diminished lateralization, we analyzed 407 the cPMI again after applying a strict procedure to exclude trails in which small eve 408 409 movements were present, based on visual inspection of the EOG traces of each trial. The results on alpha lateralization remained, as we still found a significant effect for Interval 410 (F(1,54) = 11.838, p = 0.001), Interval X Age-group (F(1,54) = 25.399, p = 0.000), and Age-411 group (F(1,54) = 18.327, p = 0.000). Thus, eye-movements could not explain the diminished 412 lateralization during the retention interval in older adults. 413

415 *Raw and baselined alpha power*

The modulation index does not provide any information on whether the lack of 416 modulation in older adults was due to alpha power being equally high or equally low in both 417 418 conditions. To tease apart the mechanisms underlying the modulation we first investigated the absolute levels of alpha power. After log-transforming the time-frequency data, cue and 419 retention interval values were combined and averaged per individual, and averaged over both 420 sensor clusters (Figure 4C). An independent samples t-test on the resulting average (log-421 transformed) alpha power values revealed that older adults (M = -27.03, SD = 0.32) showed 422 significantly lower alpha power (t(57) = 3.04, p = 0.004) than younger adults (M = -26.77, SD 423 = 0.33). Furthermore, we were able to replicate (Figure 4C) recent findings by Voytek et al. 424 425 (2015), who found that older adults have significantly flatter 1/f-noise spectra (t(57) = -3.97, p = 0.000). This could indicate more spontaneous (and thus less synchronized) high 426 frequency activity, pointing at deficiencies in the regulation of high frequency activity by 427 lower frequency oscillations such as alpha (Canolty et al. 2006; Jensen and Colgin 2007; 428 429 Bastos et al. 2015; Voytek et al. 2015; Lowet et al. 2016). Next we investigated the development of alpha power from a baseline through the cue 430 and retention intervals. Because alpha power developed differently depending on the 431 attention condition and hemisphere, those parameters were combined by labeling, per trial, 432 each hemisphere as ipsilateral or contralateral relative to the target hemifield. The log-433 434 transformed data were then sorted and averaged according to their laterality, age group, and interval. Then, from each signal a baseline (-1.75 s - -1.5 s) was subtracted, so that Figure 4A 435 and 4B show changes from baseline as a function of time. The resulting traces show that, in 436 both younger and older adults, alpha power decreased in the cue interval compared to 437 baseline. In both groups, alpha power decreased more over the hemisphere contralateral to 438 the relevant side of the array than over the ipsilateral hemisphere, leading to alpha 439

440 lateralization. In the WM retention interval, younger adults showed an initial alpha suppression caused by the onset of the sample array, followed by an ipsilateral alpha power 441 increase to baseline levels. Alpha power contralateral to the relevant side of the array 442 443 continued to be suppressed compared to the ipsilateral hemisphere. Strikingly, in older adults there was an even larger decrease in both ipsilateral and contralateral alpha power in the 444 retention interval, during which, ipsilateral and contralateral alpha power levels were both 445 reduced to a similar level. Thus, the absence of modulation in older adults during the 446 retention interval was paired with an overall bilateral decrease in alpha power. 447 448 These observations were tested with an RM-ANOVA, with Laterality (ipsilateral vs. contralateral) and Interval (cue interval vs. retention interval) as within-subject factors, and 449 450 Age-group as a between subject factor (Figure 4D). There were significant interactions between Laterality and Age-group (F(1,57) = 18.189, p = 0.000), Laterality and Interval 451 (F(1,57) = 5.139, p = 0.027), and Laterality, Interval, and Age (F(1,57) = 23.728, p = 0.000), 452 underlining the fact that ipsilateral and contralateral alpha power were affected differently by 453 454 the cue and retention intervals, and age. Paired sample t-tests confirmed that in the cue interval, both younger adults (t(24) = 5.261, p = 0.000) and older adults (t(33) = 3.522, p =455 0.001) had higher alpha power in the ipsilateral hemisphere than in the contralateral 456 hemisphere. In the retention interval this was the case for younger adults (t(24) = 5.675, p =457 0.000), but not for older adults (t(33) = 1.159, p = 0.255). Interestingly, there was no 458 459 significant difference in ipsilateral alpha power between the cue and retention intervals in younger adults (t(24) = 0.998, p = 0.328), whereas in older adults ipsilateral alpha power was 460 significantly lower in the retention than in the cue (t(33) = 5.238, p = 0.000). Contralateral 461 alpha power decreased significantly from cue to retention in both younger adults (t(24) =462 2.444, p = 0.022) and older adults (t(33) = 4.883, p = 0.000). The lack of alpha lateralization 463

observed in older adults during the retention interval was hence due mostly to a reduction inalpha power contralateral to the irrelevant side of the array.

466

467 ----- Figure 4 near here -----

468

469 *Control analyses*

The younger group was part of a pharmacological study consisting of two drug sessions 470 and one placebo session. In the current study only data from the placebo session was used. 471 However, due to the counterbalancing of drug conditions, in the younger group the placebo 472 session was not always the second session (first MEG session after the initial training and 473 474 MRI acquisition session). Therefore, some of the younger adults could be more experienced 475 with the task than participants in the older group. To test whether practice effects contributed to our findings, the main analysis on cPMI was repeated including as controls only those 476 younger adults (N=9) who received a placebo in their second session (Figure 5A). Again, an 477 RM-ANOVA, with cPMI as the dependent variable, Age-group (younger adults vs. older 478 adults) as between-subject factor, and Interval (cue interval vs. retention interval) as a within-479 subject factor, revealed similar effects to the main analysis summarized in Figure 3, including 480 roughly equal modulation of alpha lateralization in the cue interval for both age groups, and 481 different modulation in the retention interval. The analysis confirmed a significant effect for 482 Interval (F(1,41) = 4.084, p = 0.050). Post-hoc tests revealed higher cPMI in the cue interval 483 (M = 0.034, SD = 0.039) than in the retention (M = 0.031, SD = 0.055) interval. Age-group 484 also had a significant effect (F(1,41) = 47.04, p = 0.007), with younger adults (M = 0.060, SD 485 = 0.046) having higher cPMI than older adults (M = 0.025, SD = 0.029). Furthermore, the 486 interaction Age-group X Interval was significant as well (F(1,41) = 15.307, p = 0.000). 487 Independent sample t-tests within each interval revealed a significant effect for Age-group in 488

the retention interval (t(9.55) = 3.25, p = 0.009), but not in the cue interval (t(41) = -0.30, p = 0.763). Figure 5A and the associated analysis (Figure 5B) showed stronger modulation in the retention interval among younger adults than among older adults. Practice effects thus cannot explain the difference in modulation between younger and older adults.

Another possible confound was that there were on average more items on the screen for 493 younger adults than for older adults, due to the individual adjustment in load. To exclude the 494 possibility that the amount of squares in the array caused the different modulation patterns, 495 the main analysis was repeated once more, selecting only younger (N=5) and older adults 496 497 (N=24) in the memory load condition most commonly presented to older people: 3 squares per hemifield (Figure 5C). Again, the main observation was replicated (Figure 5D), with a 498 499 significant effect of Age-group (F(1,27) = 9.809, p = 0.004) and a significant interaction between Interval and Age-group (F(1,27) = 5.084, p = 0.032). In this analysis, independent 500 sample t-tests only revealed a trending effect for Age-group in the retention interval (t(4.457)) 501 = 2.358, p = 0.071), which is most likely due to the low number of younger adults in this 502 group. In the cue interval there were no significant or trending differences (t(27) = 0.761, p =503 0.453). Thus, younger and older adults exhibited similar modulations during the cue, whereas 504 during the retention interval modulation was stronger in younger adults and nearly absent in 505 older adults. 506

507

508 ------ Figure 5 near here ------

509

Finally, the male to female ratio was higher in the older group. We tested whether the effects we found could be caused by gender differences in the sample, and found that both males and females exhibited the same effect; no age-differences in cue interval lateralization and larger age-differences during the retention interval. This was summarized by the

- significant Age-group X Interval interactions for the male (F(1,31) = 38.555, p = 0.000) and
- female (F(1,24) = 5.083, p = 0.034) participants. The three-way interaction Age-group X
- 516 Interval X Gender was also significant however (F(1,55) = 6.110, p = 0.017), reflecting that
- 517 the effect of Age-group on Interval was stronger in males than in females. This may reflect an
- 518 interesting gender difference which could be explored in future research. Taken together
- these control analyses suggest that the differences in experimental procedures and gender
- 520 ratio between the two groups do not underlie our central findings.

521

Discussion

Many studies have shown that tasks which require attention to be allocated to one 522 hemifield lead to lateralized alpha power over posterior sites (e.g. Worden et al., 2000; Thut, 523 524 2006; Händel et al., 2011). Recent studies have demonstrated that this idea can be extended into the domain of WM (Sauseng et al. 2009). In addition, current data and theories suggest 525 that increased alpha power suppresses processing, while decreased alpha power facilitates 526 processing (Hanslmayer et al. 2005; Kelly et al. 2006; Rihs et al. 2007; Jensen and Mazaheri 527 2010; Händel et al. 2011). We therefore used MEG to test whether a decline in this 528 529 mechanism may underlie decreased WM performance during aging. One of the benefits of MEG over most common EEG systems is that the superior number of sensors allows for 530 greater spatial precision at the scalp level. More importantly, our experimental design 531 532 allowed us to separate the processes involved in cue-related attentional orienting from the processes involved in WM retention and WM-related attention. 533

We used a lateralized DMS task in which difficulty was individually adjusted so that all 534 participants were equally challenged and engaged. In the cue interval, the two hemispheres 535 showed the typical pattern of alpha power lateralization in both age-groups, namely that alpha 536 power was higher when target stimuli were expected in the ipsilateral hemifield, compared to 537 when they were expected in the contralateral hemifield. In the retention interval, however, the 538 539 expected alpha lateralization effect was strongly present only in the younger adults, but 540 nearly absent in the older adults. Additional analyses of the absolute power in the two hemispheres showed that this lack of modulation in older adults was paired with a bilateral 541 reduction in alpha power to the same level. Furthermore, alpha power was lower in the 542 retention interval than in the cue interval for older adults, whereas in younger adults 543 ipsilateral alpha power remained at the same level in both intervals. These results suggest that 544 the main difference between younger and older adults during the retention interval lies in a 545

deficiency to recover alpha power after an initial stimulus related drop in power in olderadults, in the hemisphere processing irrelevant stimuli.

The fact that alpha power was modulated by the same relative amount in response to a 548 directional cue in both younger and older adults, suggests that the brain relies on the same 549 mechanism to distribute attentional resources in both age groups, in line with Mok et al. 550 (2016). But what then could cause the difference in hemispheric alpha lateralization between 551 the two groups during the retention interval? One possible explanation is that there was 552 insufficient top-down drive to inhibit encoding of irrelevant stimuli at the onset of the arrays. 553 554 The exogenous onset of the sample array may have caused a redistribution of attention over the two hemifields, overriding the endogenous drive that directs attention to the target 555 location. In line with reduced top down control, we and others (Dustman et al. 1999; Voytek 556 557 et al. 2015) observed lower overall alpha power in older adults. Feedforward input may thus be more dominant in older adults. Furthermore, Sander et al. (2012b) found that the alpha 558 phase immediately after stimulus onset was more coherent across trials in older adults, 559 560 indicating that alpha processes in this age-group were more strongly affected by feedforward input. A deficit in top down drive fits with several theories in the literature, such as the early 561 inhibition deficit found in older adults by Gazzaley et al. (2008), as well as the two-562 component framework proposed by Sander et al. (2012a), which states that WM may rely on 563 the interplay of low-level feature binding processes and top-down control processes. In terms 564 565 of these theories, the deficits during retention may reflect a weakening of top-down control processes, and increased dominance of feedforward processing. However, arguing against the 566 interpretation that healthy aging coincides with a shift towards feedforward processing, we 567 found no difference in sensitivity and response bias between the age groups, as evidenced by 568 d' and criterion measures. Moreover, we found that older adults were not more likely to 569 report changes in stimuli when one occurred in the uncued array than when no change 570

571 occurred in either hemifield, as might be expected had they encoded the uncued stimuli. This suggests attentional control remained intact in healthy older adults. One reason for the lack of 572 evidence for the inhibition deficit theory in the current study could be that most studies 573 574 investigating inhibition deficit featured serially presented stimuli of varying relevance. In such non-concurrent presentations there may be no opportunity for older adults to prioritize 575 one set of stimuli over another set. Another explanation was presented by Vaden et al. (2012), 576 who also found no evidence for suppression deficits in older adults. They propose that there 577 may be a difference in task demands between the Sternberg tasks with realistic pictures and 578 579 the relatively simple displays employed in lateralization studies, which allows the older adults to suppress the irrelevant information. Furthermore, older adults did maintain a 580 reasonable level of WM performance, despite weak alpha lateralization in the retention 581 582 interval. Hence, alpha lateralization deficits in older adults no longer seemed to be an accurate electrophysiological index of WM performance deficits. 583

Despite the reduced alpha lateralization during retention, there was significant residual 584 WM performance. Interestingly, the reduction of alpha lateralization was paired with an 585 overall reduction in alpha power in both hemispheres. This finding could be seen as part of 586 the deficit in the older adults, but it could also be a correlate of a compensatory mechanism. 587 Specifically, we suggest that both hemispheres were recruited to maintain the relevant part of 588 the array in WM. A number of fMRI studies have shown that tasks which evoke lateralized 589 590 activity in younger adults evoke bilateral activity in high-functioning older adults (but lateralized activity in low-performing older adults), indicating that a shift towards bilateral 591 activity could be a compensatory strategy (Reuter-Lorenz et al. 2000; Cabeza et al. 2002; 592 Reuter-Lorenz and Cappell 2008). In line with these findings, the increase in bilateral 593 processing in our data (as reflected by the bilateral alpha power decrease) could be 594 interpreted as reflecting compensatory mechanisms. In this explanation, older adults rely on a 595

596 reconfigured retention mechanism in which alpha operates in a non-lateralized manner. The fact that this compensatory mechanism operates during retention and not during cueing 597 (where alpha lateralization was intact) is perhaps due to different but spatially overlapping 598 599 neural networks being involved in alpha lateralization when allocating attention (cueing) and WM (retention). A possible separation of mechanisms of alpha lateralization during cueing 600 and WM may underlie the observation that a compensatory strategy during aging comes into 601 existence for WM, leaving mechanisms for attentional orienting unaffected. However, it is 602 also possible that older adults switch from a lateralized to a bilateral mechanism in a task 603 604 dependent manner, without a need for different alpha generating networks for attentional orienting and WM. It is as yet unclear how the reconfigured retention mechanism operates in 605 606 older adults. Irrespective of how this reconfiguration is achieved it is noteworthy that, 607 although fairly effective, it is less effective than the processes in young adults as WM capacity (K_{span}) was reduced. 608

Our findings differ from those of Hong et al. (2015), who concluded that only younger 609 610 adults showed alpha power lateralization in anticipation of a cued stimulus. This contrasts with our data, which show a comparable alpha lateralization in younger and older age groups 611 during the cue interval, and a reduction of alpha power and lateralization during retention in 612 the older age group specifically. Thus, we suggest that the reduction in alpha lateralization 613 related to normal aging is more selective than previously thought, being only apparent during 614 615 the retention interval in our task. The difference in results between the Hong et al. (2015) study and our own may be due to differences in experimental design. In this regard, it is 616 noteworthy that in Hong et al. (2015) the target was always known to the participants, 617 whereas in our task the target was unknown to the participants during the cue interval. 618 Therefore, what they termed a cue interval in their study perhaps is more comparable to the 619 retention interval in our study, rather than to our cue interval. In this light both investigations 620

621 find that in older adults alpha power was not lateralized during WM retention. Importantly, our experimental design, which separates processes related to attentional cueing from WM-622 related processes, allowed the identification of a decline in alpha lateralization and alpha 623 power in older adults specific to WM-related operations and not to attentional spatial cueing. 624 One limitation in the current study was that because difficulty was individually 625 adjusted, we could not compare electrophysiological processes at play during high and low 626 loads, as in Sander et al. (2012b). We were also unable to demonstrate correlations between 627 individual performance and the amount of alpha power modulation as demonstrated by e.g. 628 629 Sauseng (2009). These analyses would have furthered our understanding of the performance deficits and compensatory strategies of older adults, and crucially of their underlying 630 neuronal mechanisms. However, the current design was also one of the study's strengths, as 631 632 we ensured that the task was equally difficult and engaging for younger and older adults. This was especially important considering that in some studies differences in experienced task 633 difficulty alone explained differences in brain activation (Schneider-Garces et al. 2009). 634 In conclusion, our analysis of alpha power in older and younger adults revealed 635 different mechanisms during retention in a WM task, but no differences were found in 636 response to attentional cueing without WM. In older adults, we found bilateral alpha power 637 reductions and lack of alpha lateralization during retention, which may either reflect a failure 638 to suppress distractors, or be part of a compensatory mechanism. We found that older adults 639 640 did not respond more to irrelevant items than younger adults, and that both younger and older adults showed lateralized alpha oscillations during attentional orienting. This supports our 641 tentative conclusion that mechanisms involved in attentional orienting and encoding remain 642 relatively intact during healthy aging, and that declined WM performance in our task is 643 specifically due to a reconfigured retention mechanism that is not as effective as in the young 644 adults. 645

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651	References
652 653	Bastiaansen MCM, Knösche TR. 2000. Tangential derivative mapping of axial MEG applied to event-related desynchronization research. Clin Neurophysiol. 111:1300–1305.
654 655	Bastos AM, Vezoli J, Bosman CA, Schoffelen J-M, Oostenveld R, Dowdall JR, De Weerd P, Kennedy H, Fries P. 2015. Visual Areas Exert Feedforward and Feedback Influences
656	through Distinct Frequency Channels. Neuron. 85:390–401.
657 658	Bopp KL, Verhaeghen P. 2005. Aging and verbal memory span: a meta-analysis. J Gerontol
658 659	Psychol Sci. 60B:223–233. Cabeza R, Anderson ND, Locantore JK, McIntosh AR. 2002. Aging gracefully:
660	compensatory brain activity in high-performing older adults. NeuroImage. 17:1394–
661	1402.
662	Canolty RT, Edwards E, Dalal SS, Soltani M, Nagarajan SS, Kirsch HE, Berger MS, Barbaro
663 664	NM, Knight RT. 2006. High Gamma Power Is Phase-Locked to Theta Oscillations in Human Neocortex. Science. 313:1626–1628.
665	Cowan N. 2000. The magical number 4 in short-term memory: a reconsideration of mental
666	storage capacity. Behav Brain Sci. 24:87–185.
667	De Graaf A, Deelman B. 1991. Cognitieve screening test. Lisse Swets En Zeitlinger.
668	D'Esposito M. 2007. From cognitive to neural models of working memory. Phil Trans R Soc
669	B. 362:761–772.
670	Dustman RE, Shearer DE, Emmerson RY. 1999. Life-span changes in EEG spectral
671	amplitude, amplitude variability and mean frequency. Clin Neurophysiol. 110:1399-
672	1409.
673	Gazzaley A, Clapp W, Kelley J, McEvoy K, Knight RT, D'Esposito M. 2008. Age-related
674	top-down suppression deficit in the early stages of cortical visual memory processing.
675	Proc Natl Acad Sci. 105:13122–13126.
676	Gazzaley A, Nobre AC. 2012. Top-down modulation: bridging selective attention and
677 678	working memory. Trends Cogn Sci. 16:129–135. Händel BF, Haarmeier T, Jensen O. 2011. Alpha oscillations correlate with the successful
679	inhibition of unattended stimuli. J Cogn Neurosci. 23:2494–2502.
680	Hanslmayer S, Klimesch W, Sauseng P, Gruber W, Doppelmayr M, Freunberger R,
681	Pecherstorfer T. 2005. Visual discrimination performance is related to decreased
682	alpha amplitude but increased phase locking. Neurosci Lett. 375:64–68.
683	Hautus MJ. 1995. Corrections for extreme proportions and their biasing effects on estimated
684	values ofd'. Behav Res Methods Instrum Comput. 27:46–51.
685	Hedden T, Gabrieli JDE. 2004. Insights into the ageing mind: a view from cognitive
686	neuroscience. Nat Rev Neurosci. 5:87–96.
687	Hong X, Sun J, Begson JJ, Mangun GR, Tong S. 2015. Normal aging selectively diminishes
688	alpha lateralization in visual spatial attention. NeuroImage. 106:353-363.
689	Jensen O, Colgin LL. 2007. Cross-frequency coupling between neuronal oscillations. Trends
690	Cogn Sci. 11:267–269.
691	Jensen O, Mazaheri A. 2010. Shaping functional architecture by oscillatory alpha activity:
692	gating by inhibition. Front Hum Neurosci. 4:1–8.
693	Jost K, Bryck RL, Vogel EK, Mayr U. 2011. Are old adults just like low working memory
694 605	young adults? Filtering efficiency and age differences in visual working memory.
695 696	Cereb Cortex. 21:1147–1154. Jung TP, Makeig S, Humphries C, Lee TW, McKeown MJ, Iraqui V, Sejnowski TJ. 2000.
697	Removing electroencephalographic artifacts by blind source separation.
698	Psychophysiology. 37:163–178.

- Kelly SP, Lalor EC, Reilly RB, Foxe JJ. 2006. Increases in alpha oscillatory power reflect an
 active retinotopic mechanism for distracter suppression during sustained visuospatial
 attention. J Neurophysiol. 95:3844–3851.
- Klimesch W. 2012. Alpha-band oscillations, attention, and controlled access to stored information. Trends Cogn Sci. 16:606–617.
- Lowet E, Roberts MJ, Bosman CA, Fries P, De Weerd P. 2016. Areas V1 and V2 show microsaccade-related 3–4-Hz covariation in gamma power and frequency. Eur J Neurosci. 43:1286–1296.
- Lozano-Soldevilla D, ter Huurne N, Cools R, Jensen O. 2014. GABAergic modulation of
 visual gamma and alpha oscillations and its consequences for working memory
 performance. Curr Biol. 24:2878–2887.
- Luck SJ, Vogel EK. 1997. The capacity of visual working memory for features and conjunctions. Nature. 390:279–281.
- Luck SJ, Vogel EK. 2013. Visual Working Memory Capacity: From Psychophysics and
 Neurobiology to Individual Differences. Trends Cogn Sci. 17:391–400.
- Maris E, Oostenveld R. 2007. Nonparametric statistical testing of EEG- and MEG-data. J
 Neurosci Methods. 164:177–190.
- McNab F, Zeidman P, Rutledge RB, Smittenaar P, Brown HR, Adams RA, Dolan RJ. 2015.
 Age-related changes in working memory and the ability to ignore distraction. Proc Natl Acad Sci. 112:6515–6518.
- Mok RM, Myers NE, Wallis G, Nobre AC. 2016. Behavioral and Neural Markers of Flexible
 Attention over Working Memory in Aging. Cereb Cortex. 26:1831–1842.
- Murre JMJ, Janssen SMJ, Rouw R, Meeter M. 2013. The rise and fall of immediate and delayed memory for verbal and visuospatial information from late childhood to late adulthood. Acta Psychol (Amst). 142:96–107.
- Oostenveld R, Fries P, Maris E, Schoffelen J-M. 2011. FieldTrip: Open source software for
 advanced analysis of MEG, EEG, and invasive electrophysiological data. Comput
 Intell Neurosci. 2011:1–9.
- Park DC, Lautenschlager G, Hedden T, Davidson NS, Smith AD, Smith PK. 2002. Models of
 visuospatial and verbal memory across the adult life span. Psychol Aging. 17:299–
 320.
- Pashler HH. 1988. Familiarity and visual change detection. Percept Psychophys. 44:369–378.
- Reuter-Lorenz PA, Cappell KA. 2008. Neurocognitive Aging and the Compensation
 Hypothesis. Curr Dir Psychol Sci. 17:177–182.
- Reuter-Lorenz PA, Jonides J, Smith EE, Hartley A, Miller A, Marshuetz C, Koeppe RA.
 2000. Age differences in the frontal lateralization of verbal and spatial working
 memory revealed by PET. J Cogn Neurosci. 12:174–187.
- Rihs TA, Michel CM, Thut G (2007) Mechanisms of selective inhibition in visual spatial
 attention are indexed by α-band EEG synchronization. Eur J Neurosci 25:603–610.
- Sander MC, Lindenberger U, Werkle-Bergner M. 2012. Lifespan age differences in working memory: A two-component framework. Neurosci Biobehav Rev. 36:2007–2033.
- Sander MC, Werkle-Bergner M, Lindenberger U. 2011. Binding and strategic selection in
 working memory: a lifespan dissociation. Psychol Aging. 26:612–624.
- Sander MC, Werkle-Bergner M, Lindenberger U. 2012. Amplitude modulations and inter trial phase stability of alpha-oscillations differentially reflect working memory
 constraints across the lifespan. NeuroImage. 59:646–654.
- Sauseng P, Klimesch W, Heise KF, Gruber WR, Holz E, Karim AA, Glennon M, Gerloff C,
 Birbaumer N, Hummel FC. 2009. Brain oscillatory substrates of visual short-term
 memory capacity. Curr Biol. 19:1846–1852.

- Schneider-Garces NJ, Gordon BA, Brumback-Peltz CR, Shin E, Lee Y, Sutton BP, Maclin
 EL, Gratton G, Fabiani M. 2009. Span, CRUNCH, and beyond: working memory
 capacity and the aging brain. J Cogn Neurosci. 22:655–669.
- Stolk A, Todorovic A, Schoffelen J-M, Oostenveld R. 2013. Online and offline tools for head
 movement compensation in MEG. NeuroImage. 68:39–48.
- Thut G. 2006. α-band electroencephalographic activity over occipital cortex indexes
 visuospatial attention bias and predicts visual target detection. J Neurosci. 26:9494–
 9502.
- Vaden RJ, Hutcheson NL, McCollum LA, Kentros J, Visscher KM. 2012. Older adults,
 unlike younger adults, do not modulate alpha power to suppress irrelevant information.
 NeuroImage. 63:1127–1133.
- Vogel EK, Machizawa MG. 2004. Neural activity predicts individual differences in visual
 working memory capacity. Nature. 428:748–751.
- Vogel EK, McCollough AW, Machizawa MG. 2005. Neural measures reveal individual differences in controlling access to working memory. Nature. 438:500–503.
- Vogel EK, Woodman GF, Luck SJ. 2001. Storage of features, conjunctions and objects in visual working memory. J Exp Psychol Hum Percept Perform. 27:92–114.
- Voytek B, Kramer MA, Case J, Lepage KQ, Tempesta ZR, Knight RT, Gazzaley A. 2015.
 Age-related changes in 1/f neural electrophysiological noise. J Neurosci. 35:13257– 13265.
- Worden MS, Foxe JJ, Wang N, Simpson GV. 2000. Anticipatory biasing of visuospatial
 attention indexed by retinotopically specific-band electroencephalography increases
 over occipital cortex. J Neurosci. 20:1–6.
- 771 772

Tables

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Load	Sig	Younger adults	Older adults
2		90 (±9.8) %	90 (±6.0) %
3	*	82 (±8.6) %	76 (±9.0) %
4	*	73 (±8.7) %	67 (±8.0) %
(5)		68 (±6.5) %	N/A
(6)		65 (±7.8) %	N/A

Table 1 Accuracy in session 1

Note: Load indicates number of squares in each hemifield. Asterisks indicate significant

differences in mean accuracy between younger adults and older adults. Standard deviations in

777 brackets.

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Captions to figures

Figure 1 The delayed match-to-sample task. Participants always fixated on the center 780 symbol. After an inter-trial period of 2 seconds, in which participants were free to blink, the 781 fixation cross changed into a directional cue ('<' or '>'). This cue indicated which hemifield 782 should be remembered and compared to the probe array, and which hemifield should be 783 ignored. The cue remained visible for the remainder of the trial. After the 1500 ms cue 784 interval a sample array was shown for 100 ms, consisting of multiple colored squares. 785 Participants had to retain information about the color of squares in the cued hemifield during 786 787 a 1500 ms retention interval. Finally, a probe array was shown, in which one square per hemifield could have changed color. No duplicate colors were possible. The positions of 788 789 squares never changed within a trial, but varied between trials. The number of squares per 790 hemifield was the memory load and was specific for each participant (titrated to ~75% accuracy). The memory load was fixed for the entire MEG experiment. Loads ranged from 2 791 to 6 squares across younger adults, and from 2 to 4 squares across older adults (see Results). 792 793 Participants had to report within 2 seconds whether the probed squares in the cued hemifield were identical or different from the sample array. The correct response in this example would 794 be 'different'. 795

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Figure 2 A) Grand average alpha Power Modulation Index (PMI) topographical plot.
Sensors are marked as dots, and sensors that significantly differed between attend left and
attend right conditions are marked as bold dots. Significant sensors indicated by white dots
were left out of the final analysis because there were no significant sensors that mirrored
them in the opposite hemisphere. The positive and negative sensor clusters were found by
employing a cluster-based permutation test on the grand-average cue-interval (not shown). B)
Topographical plots and time frequency representations belonging to the positive cluster

(left) and negative cluster (right) in younger adults, showing the average PMI. Topographical
plots show activity during the retention interval. Dashed boxes indicate the range of
frequencies and latencies that were averaged and included in statistical analysis. C) Identical
to B, but showing data from older adults. D) Average alpha PMI for both age groups. Dashed
vertical lines indicate different epochs within a trial. Shaded areas represent standard error of
the mean.

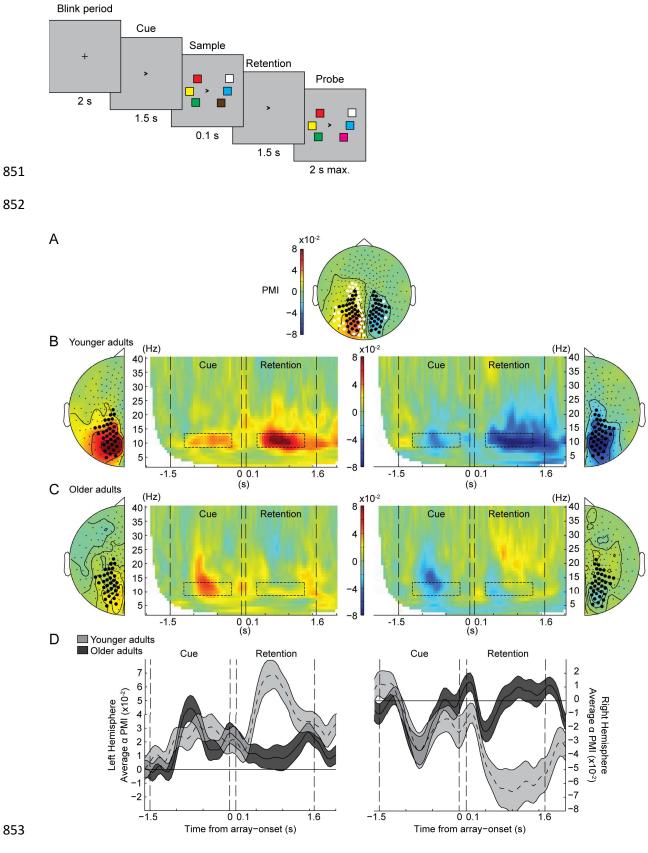
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Figure 3 The combined Power Modulation Index (cPMI) in the alpha band (8-14 Hz), for younger and older adults per interval, calculated by subtracting right hemisphere alpha PMI from left hemisphere alpha PMI. There was no difference between older and younger adults in cue interval cPMI, but in the retention interval there was a significant difference. The effect of age is also different in the two intervals, indicated by a significant interaction between age and interval. Asterisks indicate significance (*** p = 0.000; n.s. = not significant).

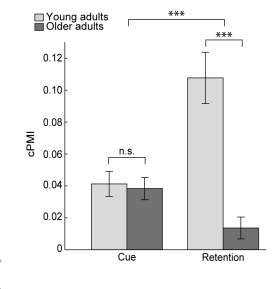
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Figure 4 A) Log-ratio between alpha power and baseline (in dB), averaged over 819 younger adults. Darker colors indicate ipsilateral alpha power, lighter colors indicate 820 contralateral alpha power. B) Like A, but averaged over older adults. C) Log-transformed 821 power spectrum for younger (blue) and older (red) adults, averaged over cue and retention 822 intervals. Dashed lines represent linear fits of 1/f noise (see Voytek et al., 2015). The shaded 823 area indicates the alpha band. D) Log-transformed alpha power, relative to baseline, averaged 824 separately over the cue and retention intervals. Significance of paired t-tests is indicated by 825 asterisks (*** p = 0.000). 826

828	Figure 5 Mean alpha Power Modulation Index (PMI) comparisons between older adults and
829	younger adults. A) Mean alpha PMI for older adults and younger adults that were recorded in
830	the second session (rather than session 3 or 4), in the same format as Figure 2D. Shaded areas
831	show standard error of the mean. B) Mean alpha combined PMI for young and old adults
832	from data recorded in the second session, in the same format as Figure 3. C) Mean alpha PMI
833	for older adults and younger adults in conditions where there were always 3 squares per
834	hemifield on the screen. D) Mean alpha combined PMI for young and old adults from data
835	recorded when there were 3 squares per hemifield on the screen. Note that there are still only
836	small differences between age groups in the cue interval (-1.5 s $-$ 0 s) and large differences in
837	the retention interval ($0.1 \text{ s} - 1.6 \text{ s}$).
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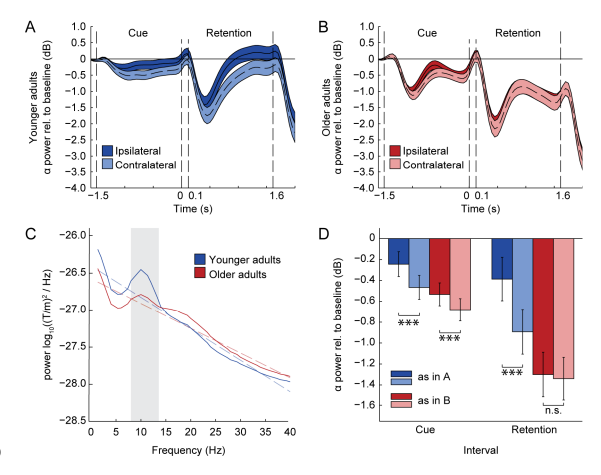


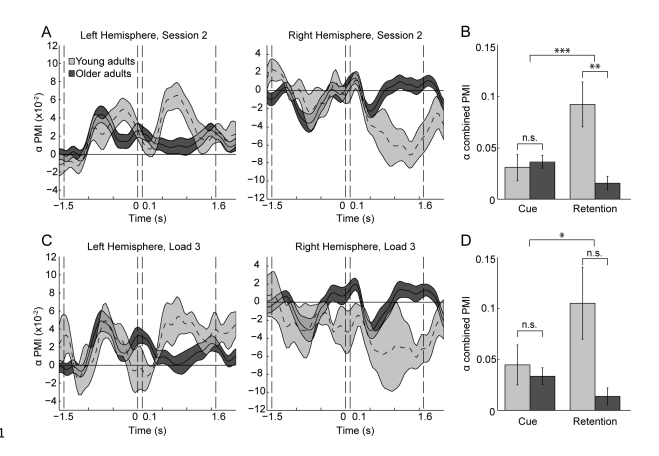














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